Entry name	CLPP_ECOLI				
Primary accession number	P19245				
Secondary accession numbers	None				
Entered in Swiss-Prot in	Release 16, November 1	.990			
Sequence was last modified in	Release 16, November 1	.990			
Annotations were last modified in	Release 41, February 20	003			
Name and origin of the protein					
Protein name	ATP-dependent Clp prosubunit	otease proteolytic			
Synonyms	EC 3.4.21.92				
	Endopeptidase Clp				
	Caseinolytic protease				
	Protease Ti				
	Heat shock protein F2	1.5			
Gene name	CLPP or LOPP or <u>B0437</u> <u>Z0542</u> or ECS0491 or				
From	Escherichia coli	[TaxID: <u>562]</u>			
	Escherichia coli 06	[TaxID: <u>217992]</u>			
	Escherichia coli 0157:H7	[TaxID: <u>83334]</u>			
	Shigella flexneri	[TaxID: <u>623]</u>			
Taxonomy	Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Escherichia.				

#### References

[1] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli;

MEDLINE=90324245; PubMed=2197275; [<u>NCBI</u>, <u>ExPASy</u>, <u>EBI</u>, <u>Israel</u>, Japan]

Maurizi M.R., Clark W.P., Katayama Y., Rudikoff S., Pumphrey J., Bowers B., Gottesman S.;

"Sequence and structure of Clp P, the proteolytic component of the

ATP-dependent Clp protease of Escherichia coli."; J. Biol. Chem. 265:12536-12545(1990).

[2] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli;

**STRAIN**=K12 / MG1655;

MEDLINE=97426617; PubMed=9278503; [<u>NCBI</u>, <u>ExPASy</u>, <u>EBI</u>, <u>Israel</u>, <u>Japan</u>]

Blattner F.R., Plunkett G. III, Bloch C.A., Perna N.T., Burland V., Riley M., Collado-Vides J., Glasner J.D., Rode C.K., Mayhew G.F., Gregor J., Davis N.W., Kirkpatrick H.A., Goeden M.A., Rose D.J., Mau B., Shao Y.; "The complete genome sequence of Escherichia coli K-12."; Science 277:1453-1474(1997).

[3] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli;

Roberts D., Allen E., Araujo R., Aparicio A., Chung E., Davis K., Duncan M., Federspiel N., Hyman R., Kalman S., Komp C., Kurdi O., Lew H., Lin D., Namath A., Oefner P., Schramm S., Davis R.W.;
Submitted (JAN-1997) to the EMBL/GenBank/DDBJ databases.

[4] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli;

**STRAIN**=06:H1 / CFT073 / ATCC 700928;

MEDLINE=22388234; PubMed=12471157; [NCBI, ExPASy, EBI, Israel, Japan]

Welch R.A., Burland V., Plunkett G. III, Redford P., Roesch P., Rasko D., Buckles E.L., Liou S.-R., Boutin A., Hackett J., Stroud D., Mayhew G.F., Rose D.J., Zhou S., Schwartz D.C., Perna N.T., Mobley H.L.T., Donnenberg M.S., Blattner F.R.;

"Extensive mosaic structure revealed by the complete genome sequence of uropathogenic Escherichia coli.";

<u>Proc. Natl. Acad. Sci. U.S.A. 99:17020-17024(2002)</u>.

[5] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli:

**STRAIN**=0157:H7 / EDL933 / ATCC 700927;

MEDLINE=21074935; PubMed=11206551; [<u>NCBI</u>, <u>ExPASy, EBI, Israel,</u> Japan] Perna N.T., Plunkett G. III, Burland V., Mau B., Glasner J.D., Rose D.J., Mayhew G.F., Evans P.S., Gregor J., Kirkpatrick H.A., Posfai G., Hackett J., Klink S., Boutin A., Shao Y., Miller L., Grotbeck E.J., Davis N.W., Lim A., Dimalanta E.T., Potamousis K., Apodaca J., Anantharaman T.S., Lin J., Yen G., Schwartz D.C., Welch R.A., Blattner F.R.; "Genome sequence of enterohaemorrhagic Escherichia coli 0157:H7."; Nature 409:529-533(2001).

[6] SEQUENCE FROM NUCLEIC ACID.

SPECIES=E.coli;

STRAIN=0157:H7 / RIMD 0509952:

MEDLINE=21156231; PubMed=11258796; [NCBI, ExPASy, EBI, Israel, Japan]

Hayashi T., Makino K., Ohnishi M., Kurokawa K., Ishii K., Yokoyama K., Han C.-G., Ohtsubo E., Nakayama K., Murata T., Tanaka M., Tobe T., Iida T., Takami H., Honda T., Sasakawa C., Ogasawara N., Yasunaga T., Kuhara S., Shiba T., Hattori M., Shinagawa H.;

"Complete genome sequence of enterohemorrhagic Escherichia coli O157:H7 and genomic comparison with a laboratory strain K-12."; DNA Res. 8:11-22(2001).

[7] SEQUENCE FROM NUCLEIC ACID.

SPECIES=S.flexneri;

STRAIN=301 / Serotype 2a;

MEDLINE=22272406; PubMed=12384590; [<u>NCBI, ExPASy, EBI, Israel, Japan]</u>

Jin Q., Yuan Z., Xu J., Wang Y., Shen Y., Lu W., Wang J., Liu H., Yang J., Yang F., Zhang X., Zhang J., Yang G., Wu H., Qu D., Dong J., Sun L., Xue Y., Zhao A., Gao Y., Zhu J., Kan B., Ding K., Chen S., Cheng H., Yao Z., He B., Chen R., Ma D., Qiang B., Wen Y., Hou Y., Yu J.;

"Genome sequence of Shigella flexneri 2a: insights into pathogenicity through comparison with genomes of Escherichia coli K12 and O157."; Nucleic Acids Res. 30:4432-4441(2002).

[8] IDENTIFICATION AS A HEAT SHOCK PROTEIN.

SPECIES=E.coli;

MEDLINE=91008981; PubMed=2211522; <u>[NCBI, ExPASy, EBI, Israel,</u> Japan] Kroh H.E., Simon L.D.;

"The ClpP component of Clp protease is the sigma 32-dependent heat shock protein F21.5.";

<u>J. Bacteriol. 172:6026-6034(1990)</u>.

[9] CHARACTERIZATION.

SPECIES=E.coli;

MEDLINE=94012667; PubMed=8407953; [<u>NCBI</u>, <u>ExPASy</u>, <u>EBI</u>, <u>Israel</u>, <u>Jap</u>an]

Arribas J., Castano J.G.;

"A comparative study of the chymotrypsin-like activity of the rat liver multicatalytic proteinase and the ClpP from Escherichia coli.";

J. Biol. Chem. 268:21165-21171(1993).

[10]X-RAY CRYSTALLOGRAPHY.

SPECIES=E.coli;

MEDLINE=96428678; PubMed=8831780; [NCBI, ExPASy, EBI, Israel, Japan]

Shin D.H., Lee C.S., Chung C.H., Suh S.W.;

"Molecular symmetry of the ClpP component of the ATP-dependent Clp protease, an Escherichia coli homolog of 20 S proteasome.";

<u>J. Mol. Biol. 262:71-76(1996)</u>.

[11] X-RAY CRYSTALLOGRAPHY (2.3 ANGSTROMS) OF 25-207.

SPECIES=E.coli;

MEDLINE=98050920; PubMed=9390554; [NCBI, ExPASy, EBI, Israel, Japan]

Wang J., Hartling J.A., Flanagan J.M.;

"The structure of ClpP at 2.3-A resolution suggests a model for ATP-dependent proteolysis.";

Cell 91:447-456(1997).

### Comments

FUNCTION: Cleaves peptides in various proteins in a process that requires ATP hydrolysis. Has a chymotrypsin-like activity. Plays a major role in the degradation of misfolded proteins. May play the role of a master protease which is attracted to different substrates by different specificity factors such as clpA or clpX.

CATALYTIC ACTIVITY: Hydrolysis of proteins to small peptides in the presence of ATP and magnesium. Alpha-casein is the usual test substrate. In the absence of ATP, only oligopeptides shorter than five residues are cleaved (such as succinyl-Leu-Tyr-|-NHMEC; and Leu-Tyr-Leu-|-Tyr-Trp, in which the cleavage of the -Tyr-|-Leu- and -Tyr-|-Trp- bond also occurs).

SUBUNIT: 14 CLPP SUBUNITS ASSEMBLE INTO A DISK-LIKE STRUCTURE WITH A CENTRAL CAVITY, RESEMBLING THE STRUCTURE OF EUKARYOTIC PROTEASOMES. IN THE PRESENCE OF ATP, CLPA OR CLPX SUBUNITS INTERACT WITH THE CLPP STRUCTURE TO FORM A 750 kDa COMPLEX THAT EXHIBITS ATP-DEPENDENT PROTEOLYTIC ACTIVITY.

SUBCELLULAR LOCATION: Cytoplasmic.

INDUCTION: By heat shock.

SIMILARITY: BELONGS TO PEPTIDASE FAMILY S14.

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#### Cross-references

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## Keywords

Hydrolase; Serine protease; Heat shock; 3D-structure; Complete proteome.

## Features

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Feature table viewer

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BLAST submission on ExPASy/SIB or at NCBI (USA)



Sequence analysis tools: <u>ProtParam</u>, <u>ProtScale</u>, <u>Compute pI/Mw</u>, <u>PeptideMass</u>, <u>PeptideCutter</u>, <u>Dotlet</u> (Java)



ScanProsite, MotifScan



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9 of 9

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General information about the entry

Entry name	CLPP_YEREN
Primary accession number	Q60107
Secondary accession numbers	None
Entered in Swiss-Prot in	Release 35, November 1997
Sequence was last modified in	Release 35, November 1997
Annotations were last modified in	Release 41, February 2003
Name and origin of the protein	

Name and origin of the protein	
Protein name	ATP-dependent Clp protease proteolytic subunit
Synonyms	EC <u>3.4.21.92</u> Endopeptidase Clp
Gene name	CLPP
From	<u>Yersinia enterocolitica</u> [TaxID: <u>630</u> ]
Taxonomy	Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Yersinia.

#### References

[1]|SEQUENCE FROM NUCLEIC ACID.

STRAIN=8081C / Serotype O:8;

MEDLINE=98043541; PubMed=9383193; [<u>NCBI, ExPASy, EBI, Israel,</u> Japan]

<u>Oupunj</u> Dadansan K.T., Canta

Pederson K.J., Carlson S., Pierson D.E.;

"The ClpP protein, a subunit of the Clp protease, modulates ail gene expression in Yersinia enterocolitica.";

Mol. Microbiol. 26:99-107(1997).

### Comments

FUNCTION Cleaves peptides in various proteins in a process that requires ATP hydrolysis. Has a chymotrypsin-like activity. Plays a major role in the degradation of misfolded proteins (By similarity).

CATALYTIC ACTIVITY: Hydrolysis of proteins to small peptides in the presence of ATP and magnesium. Alpha-casein is the usual test substrate. In the absence of ATP, only oligopeptides shorter than five residues are

cleaved (such as succinyl-Leu-Tyr-|-NHMEC; and Leu-Tyr-Leu-|-Tyr-Trp, in which the cleavage of the -Tyr-|-Leu- and -Tyr-|-Trp- bond also occurs).

SUBCELLULAR LOCATION: Cytoplasmic (By similarity). SIMILARITY: BELONGS TO PEPTIDASE FAMILY S14.

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Cross-referen	nces
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HSSP	P19245; 1TYF. [HSSP ENTRY / PDB]
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HAMAP	MF_00444; -; 1.
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Pfam	<u>PF00574</u> ; CLP_protease; 1.
PRINTS	PROO127; CLPPROTEASEP.
TIGRFAMs	TIGR00493; clpP; 1.
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ProDom	[Domain structure / List of seq. sharing at least 1 domain].
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SWISS-2DPA	GE <u>Get region on 2D PAGE</u> .

Keywords

### <u>Hydrolase; Serine protease.</u>

#### **Features**

ACT SITE

Key From To Length Description

ACT\_SITE 111 111 BY SIMILARITY.

136

BY SIMILARITY.



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#### Sequence information

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View entry in raw text format (no links)
Report form for errors/updates in this Swiss-Prot entry

BLAST

BLAST submission on ExPASy/SIB or at NCBI (USA)



Sequence analysis tools: <u>ProtParam</u>, <u>ProtScale</u>, <u>Compute pI/Mw</u>, <u>PeptideMass</u>, <u>PeptideCutter</u>, Dotlet (Java)



ScanProsite, MotifScan



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General information about the entry